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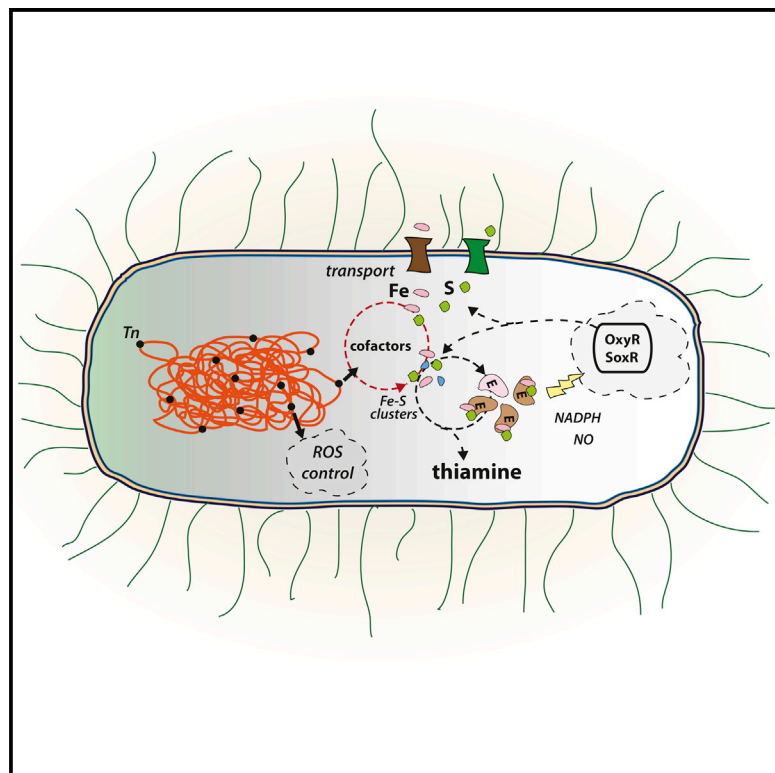
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# Cell Reports

## Genetic-Metabolic Coupling for Targeted Metabolic Engineering

### Graphical Abstract



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### In Brief

Cardinale et al. investigate the specific impact on the cell of synthetic metabolic pathways. By linking the growth of bar-coded mutants with end-product biosensing in *E. coli*, the authors identify genetic targets in the cellular Fe-S cluster biosynthesis specifically affecting the production of recombinant thiamine.

### Highlights

- Linking mutant growth to metabolite output finds product-specific genetic targets
- Genetic-metabolic coupling uncovers roles for uncharacterized genes
- Fe and S assimilation and storage differentially affect vitamin TPP production
- TPP biosynthesis triggers alternative Fe-S cluster regulation mediated by OxyR-Fur



# Genetic-Metabolic Coupling for Targeted Metabolic Engineering

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## SUMMARY

Production of chemicals in microbes often employs potent biosynthetic enzymes, which can interact with the microbial native metabolism to affect cell fitness and product yield. However, production optimization largely relies on data collected from wild-type strains in the absence of metabolic perturbations, thus limiting their relevance to specific conditions. Here, we address this issue by coupling cell fitness to the production of thiamine diphosphate in *Escherichia coli* using a synthetic RNA biosensor. We use this strategy to interrogate a library of transposon mutants and elucidate the native gene network influencing both cell fitness and thiamine production. Ultimately, we identify effectors of the OxyR-Fur stress response that limit thiamine biosynthesis via alternative regulation of iron storage and Fe-S cluster inclusion in enzymes. This study presents a new approach for the reliable high-throughput identification of genetic targets of both known and unknown function that are directly relevant to a specific biosynthetic process.

## INTRODUCTION

A common problem in the field of biotechnology is the unpredictability inherent in engineering complex biological systems. The use of a whole-system approach can prevent some of the obstacles encountered in bioengineering, including toxicity due to metabolite overproduction (Fletcher et al., 2016), metabolic bottlenecks (Lechner et al., 2016), and low product titers (Otero and Nielsen, 2010). Systems metabolic engineering relies on the generation and analysis of large datasets to identify key genetic components that contribute to high-yield production phenotypes. However, this methodology often assumes an unperturbed system, and our capacity to elucidate all metabolic and regulatory perturbations affecting the production of a given metabolite is limited.

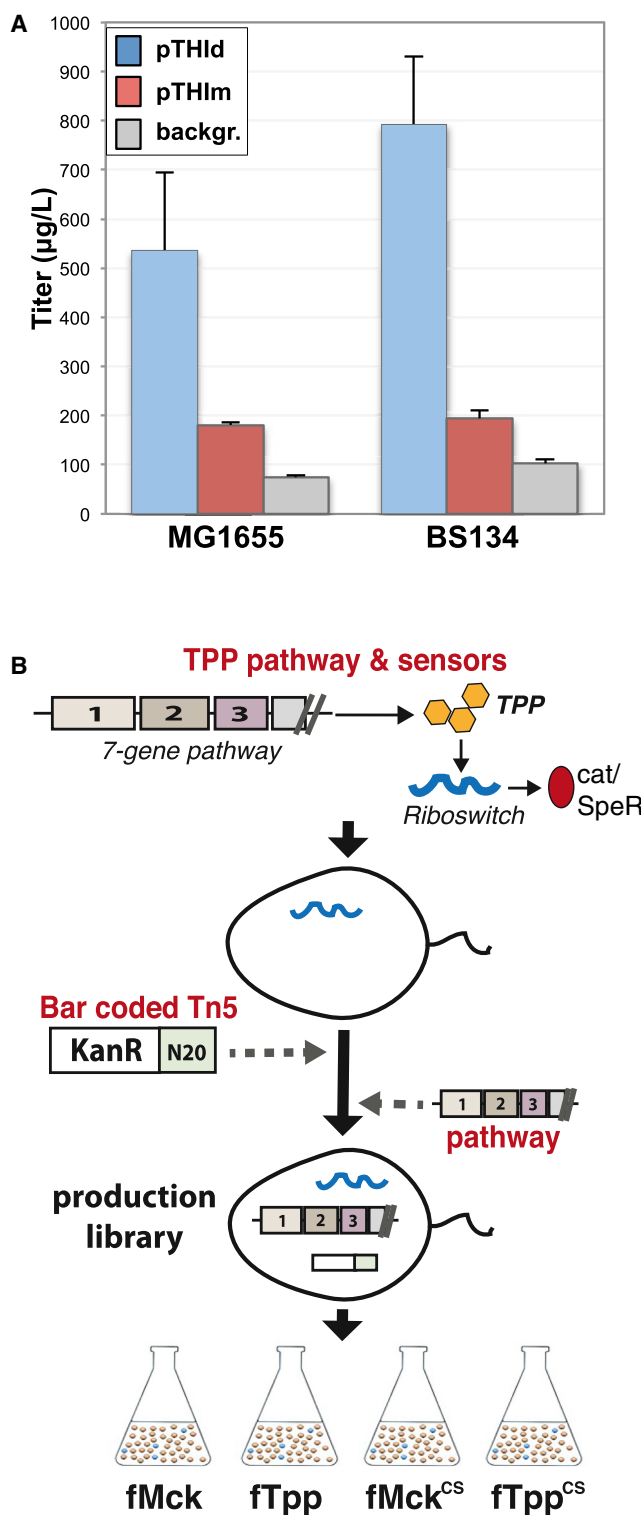
The impact of engineering a cellular system to produce a primary metabolite can be substantial. The biosynthetic pathways of primary metabolites often utilize specialized enzymes of high catalytic rate and substrate affinity; these attributes support high flux through the pathway, even at low enzyme

concentrations (Nam et al., 2012). When these enzymes are overexpressed, the supply of energy, cofactors, and carbon building blocks quickly become limiting and can lead to a strong metabolic burden on the cell. Indeed, metabolic burden can lead to low cell density and reduced titers during the production of recombinant vitamins, such as cobalamin (vitamin B<sub>12</sub>) (Biedendieck et al., 2010) or riboflavin (vitamin B<sub>2</sub>) (Lin et al., 2014). Furthermore, central cell metabolism, which provides building blocks for vitamin biosynthesis, is subject to complex regulation that can affect pathway optimization (Lin et al., 2014).

The effect of producing a metabolite is strongly dependent on the individual metabolite and how it affects cell metabolism and its regulation. Understanding the relevance of this context dependency at the genomic level remains a challenge in the field of biotechnology. New approaches enabling high-throughput characterization of fitness and pathway yields are key to achieving this goal. Tagged transposon mutagenesis (Oh et al., 2010) and synthetic RNA-based biosensors (Townshend et al., 2015) are genetic tools that have been applied to study the condition-dependent contributions of genes to cell growth and synthetic pathway output.

Here, we present a methodology for characterizing the metabolic perturbations that are a direct result of the overproduction of a primary metabolite in a bacterial cell. Genetic-metabolic coupling is based on the concurrent expression of a synthetic metabolic pathway and an end-product biosensor in a population of bar-coded single-gene transposon insertion mutants. Mutant fitness data quantified by deep sequencing are used to identify which genes provide growth advantage exclusively upon selection for the end product. We use genetic-metabolic coupling to assess the influence of *Escherichia coli* cell functions during the biosynthesis of a derivative of vitamin B<sub>1</sub>: thiamine diphosphate (TPP).

We found that the OxyR-Fur-mediated regulation of Fe-S cluster formation strongly affects TPP output. Ultimately, genetic-metabolic coupling allowed the identification of a small set of genes, several of which of uncharacterized or predicted function, with strong population-wide fitness phenotypes from over 2,000 tested genes. When introducing specific knockouts in a clean genetic background, the identified genes resulted in the predicted changes to the TPP titers, with different production plasmids and in different host strains. We finally demonstrate reverse tuning of several identified genetic targets with multi-copy plasmids, further supporting the broad application of genetic-metabolic coupling for metabolic engineering.



**Figure 1. Genetic-Metabolic Coupling Design**

(A) Quantification of extracellular TPP titer with pTHld and pTHlm (blue and red, respectively) compared to a control (no plasmid, gray) in *E. coli* MG1655 and BS134 strains ( $n = 4-6$ ). Error bars indicate SD.

(B) Graphical scheme of the experimental design. A wild-type *E. coli* strain carrying the dual riboswitch (pTPP\_Bios) is subject to mutagenesis via a

## RESULTS

### Coupling Population Genetics to Thiamine Metabolic Engineering

To create a TPP-overproducing strain for our analysis, we assembled the TPP biosynthetic genes on a plasmid. The genes were arranged in two operons that encoded the two different cellular branches required for the biosynthetic pathway. Four genes are involved in the synthesis of thiazole monophosphate (THZ-P) from L-cysteine, L-tyrosine, and deoxy-D-xylulose-5-phosphate (DXP): *thiFSGH*; *thiC*, for the synthesis of 4-amino-5-hydroxymethyl-2-methylpyrimidine phosphate (HMP-P) from 5-aminoimidazole riboside (AIR); *thiE* for coupling THZ-P to HMP-PP; and either *thiD*, the kinase that phosphorylates HMP-P to HMP-PP (plasmid pTHld), or *thiM* (plasmid pTHlm), a salvage enzyme that phosphorylates THZ. In the latter case, the strain relies on the native chromosomal *thiD* gene for HMP-P phosphorylation (details are provided in Figures S1A and S1B). The synthesis of TPP relies on several cellular biosynthetic pathways for substrates and, in particular, purine nucleotide biosynthesis for AIR, cysteine biosynthesis, and S-adenosylmethionine (SAM) and NADPH as cofactors.

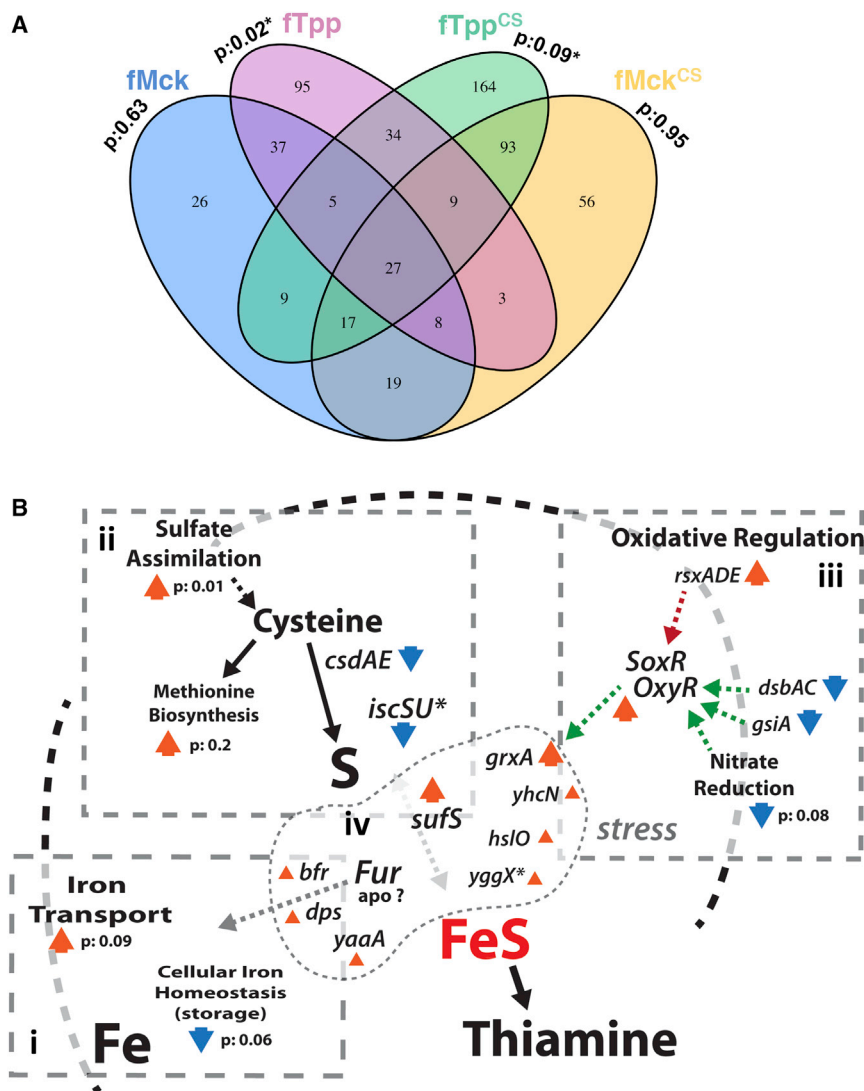
To quantitatively relay the intracellular TPP concentration at the single-cell level, we used a TPP biosensor (plasmid pTPP\_Bios) based on an engineered TPP riboswitch (Genee et al., 2016). This riboswitch is used to activate the expression of two genes conferring resistance to the antibiotics chloramphenicol (cat) and spectinomycin (SpeR). This dual-selection design allows for the reliable selection of high TPP producers using both the pTHlm and pTHld plasmids with very low false-positive rates (Genee et al., 2016). The extracellular titers of de novo-produced TPP in *E. coli* MG1655:: $\Delta$ tdpA $\Delta$ thiM (BS134), which carries deletions of the thiamine salvage and import pathways, were  $\sim 200$   $\mu$ g/L and  $\sim 800$   $\mu$ g/L for pTHlm and pTHld, respectively (Figure 1A). The overexpression of thiamine biosynthetic genes reduced the growth rates by 10% and 35% for pTHlm and pTHld, respectively.

To understand the cellular components impacting the *E. coli* TPP biosynthetic capacity, we used whole-genome transposon (Tn) mutagenesis to construct libraries of *E. coli* MG1655 Tn5 mutant strains harboring the thiamine selection system (pTPP\_Bios) and either the plasmid carrying both de novo and salvage biosynthetic genes (pTHlm) or the plasmid without a biosynthetic pathway, constructing the respective libraries ITPp and IMck (Figure 1B). Insertions were evenly distributed across the genome and through the lengths of open reading frames (Figures S1C and S1D) (RefSeq: NC\_000913.3). To characterize the impact of Tn insertions within various *E. coli* genes on the intracellular excess of TPP, we subjected the IMck and ITPp mutant libraries to a competitive fitness assay for 16 hr in shake flasks, obtaining robust fitness estimates for 1,857 genes in the IMck library and 2,054 genes in the

tagged Tn5 Tn. Following library characterization (transposon-sequencing [TnSeq]), the TPP production plasmid (pTHlm) or the backbone plasmid (pMck) is electroporated in the pool of mutants to obtain, respectively, the ITPp and IMck libraries. These libraries are finally used for fitness assays in the absence or presence (<sup>CS</sup>) of selection for TPP production.







**Figure 3. Diverse Effects of Fe-S Cluster Metabolism**

(A) Venn diagram of sets of genes with significant fitness for each cultivation condition (p: the p value of Fe-S metabolism enrichment for each condition found via mining gene annotations, Asterisk indicates significance, bootstrap n = 10,000; null hypothesis: enrichments can occur with the same probability in a random set of genes of equal size). (B) Map of the Fe-S biogenesis and regulatory networks. Four sub-networks are represented: iron supply (i), sulfur supply (ii), redox regulation of oxyR-soxR regulons (iii), and Fe-S metabolism (iv) (orange and blue arrowheads indicate improved and reduced fitness; small size: 1–2 SD from mean; big size: > 2 SD from mean). See also [Supplemental Experimental Procedures](#) and [Table S2](#).

levels (Figure 2C,  $p < 0.02$ , red bars). These effects were more significant at 5-fold higher compound concentrations, (Figure 2C, blue bars), indicating that free iron and Fe-S clusters are limiting during TPP production and that the removal of ROS has an effect similar to that of the OxyR knockouts and regulon downregulation.

### Elucidation of the Fe-S Cluster Biogenesis Network

We used a machine-learning algorithm to mine RefSeq gene descriptions (Tatusova et al., 2014) and quantified for each condition the effects of knocking out genes with biological functions related to iron, sulfur, cysteine, tyrosine, and ribose, which are cofactors or substrates involved in TPP biosynthesis (Chatterjee et al., 2008; Kriek et al., 2007). We also looked at cellular redox regulation, which

availability (*csdE*, *cysE*, and *ccmC* mutations) presented reduced fitness (Figure 2B, bottom), while gene knockouts that were involved in sulfur assimilation, major redox regulators OxyR and SoxR, and hsd DNA restriction genes resulted in a positive fitness shift (Figure 2B, top). Indeed, the pTHld pathway was more stable in a  $\Delta$ hsdMR knockout compared to wild-type (Figure S3), possibly suggesting direct cleavage by this endonuclease.

SAM turnover during TPP biosynthesis requires extensive NADPH reductive power (Chen et al., 2015). NADPH can increase cellular oxidation, impair Fe-S cluster formation, and cause accumulation of hydrogen peroxide and reactive oxygen species (ROS), all tightly controlled by OxyR-SoxR regulons (Imlay, 2015). We tested the effect on TPP titer from limiting Fe-S clusters and the activation of OxyR-SoxR regulons. Sequestration of intracellular unbound iron by iron chelators o-phenanthroline and 2,2'-dipyridyl (Imlay et al., 1988) led to a small but significant reduction in TPP titer already at low concentrations (~25  $\mu$ M), whereas thiourea, a ROS scavenger, showed near-normal TPP

is believed to influence cellular vitamin metabolism (Dougherty and Downs, 2006). We identified 108 genes in the *E. coli* genome related to these functions, among which, 57 genes (~53%) showed significant fitness shifts during selection for TPP production: 35 were associated with cofactor supply, and 25 were associated with cellular redox regulation (fTpp<sup>CS</sup> and fTpp;  $p < 0.1$  compared to a random set by random sampling with replacement or bootstrapping; Figure 3A). The presence of a significant number of genes involved in redox regulation suggests that oxidative stress can be relevant during TPP production.

To extend our analysis beyond a specific enzyme or transporter, we quantified the roles of the production phenotypes of parent GO classes (Keseler et al., 2009) associated with genes with extreme fitness phenotypes. We found that mutants of “cellular iron ion homeostasis” (GO: 006879) genes had a significant low-fitness phenotype during selection for TPP ( $p < 0.1$ ; Table S2); however, mutations in genes involved in iron, siderophores, and ferric-enterobactin transport (GO: 00015682,

0015684, 0015685, and 015891) were associated with significantly higher fitness ( $p < 0.1$ ; Figure 3Bi; Figures S4A and S4B).

In bacteria, sulfur is mainly assimilated in the cell in a reduced form, as hydrogen sulfide from sulfate, and is incorporated into the amino acids cysteine and methionine. We found that the average fitness of mutants in genes involved in “sulfate assimilation” (GO: 0000103) was significantly increased ( $p < 0.01$ ) during selection for TPP (Figure 3Bii; Figure S4C). We also found that impairment of genes related to “methionine biosynthetic process” (GO:0009086), but not cysteine biosynthesis ( $\Delta F$ :  $\Delta cysE$ ,  $-1.5$ ), led to increased fitness ( $p < 0.1$ ) (Figures S4D and S4E). In *E. coli*, the sulfur found in Fe-S clusters is derived from cytosolic L-cysteine mainly through the enzyme *IscS* (cysteine desulfurase) and the conversion of L-cysteine to L-alanine. We could not obtain fitness data for members of the ISC complex, likely because of its essentiality, but the deletion of the second sulfur-accepting complex—the CSD sulfur transfer system—showed reduced fitness upon selection for TPP ( $\Delta F$ :  $\Delta csdA$ ,  $-0.9$ ;  $\Delta csdE$ ,  $-3.4$ ) (Figure 3Bii). Surprisingly, we found that mutants in the SUF system for iron-sulfur cluster assembly that is used under iron starvation or oxidative stress conditions (Outten et al., 2004) exhibited higher fitness ( $\Delta F$ ,  $\Delta sufS$ :  $+0.4$ ,  $\Delta sufA$ :  $+0.7$ ) (Figure 3Bii).

The process of inclusion of Fe-S clusters in enzymes in *E. coli* is highly sensitive to the environment and its redox state (Crack et al., 2014; Ding et al., 2005). Tn insertions in *soxR* and *oxyR* both showed higher fitness during selection for TPP ( $\Delta F$ , respectively,  $+1.0$  and  $+1.13$ ). In addition to the *suf* operon, we found that insertions in other components of this response pathway had similar effects: the iron chelators *yaaA* and *dps* ( $\Delta F$ :  $+1.4$  and  $+0.6$ , respectively); *yhcN* ( $\Delta F$ :  $+0.6$ ), which is induced during peroxide stress (Lee et al., 2010); and the chaperone Hsp33 ( $\Delta F$ :  $\Delta hslO$   $+0.4$ ). During oxidative stress, both OxyR and SoxR upregulate the *fur* regulon that is regarded to play a key role in maintaining the stability of cellular iron levels (Seo et al., 2014). Ferritin and enterobactin have, respectively, key iron storage and iron uptake functions in *E. coli*. Bacterioferritin, whose expression is controlled by Fur via the small regulatory RNA RyhB, is important for sequestering iron, thus limiting cellular oxidative damage caused by Fenton chemistry (Bou-Abdallah et al., 2002). We found that, while Tn insertions in iron-storage ferritin and enterochelin esterase, which releases iron from ferric enterobactin, resulted in fitness defects ( $\Delta F$ :  $\Delta ftnAB$ ,  $\sim -2.0$ ;  $\Delta fes$ ,  $-3.3$ ), the lack of bacterioferritin had a positive effect on fitness during selection for TPP ( $\Delta F$ :  $\Delta bfr$ ,  $+1.0$ ) (Figure 3Biv).

We found that the control of OxyR-SoxR activation also had a significant effect. OxyR is activated by oxidation, likely by oxidized glutathione, for which just a Cys<sup>199</sup>-Cys<sup>208</sup> disulfide bond appears to be sufficient (Georgiou, 2002). We found that the impairment of glutathione import to the cytosol and *E. coli* disulfide bond system enzymes all result in negative fitness ( $\Delta F$ :  $\Delta gsaA$ ,  $-1.4$ ;  $\Delta dsbA$ ,  $-2.0$ ;  $\Delta dsbC$ ,  $-1.7$ ). In contrast, the lack of *soxR* inactivation by the RSX membrane reducing system (Koo et al., 2003) showed positive growth phenotypes ( $\Delta F$ :  $\Delta rsxA$ ,  $+1$ ;  $\Delta rsxD$ ,  $+0.8$ ;  $\Delta rsxE$ ,  $+0.6$ ) (Figure 3Biii).

Overall, our data suggest that TPP overproduction likely causes oxidative stress, as previously reported (Kriek et al., 2007), and probably iron deprivation, resulting in an activation

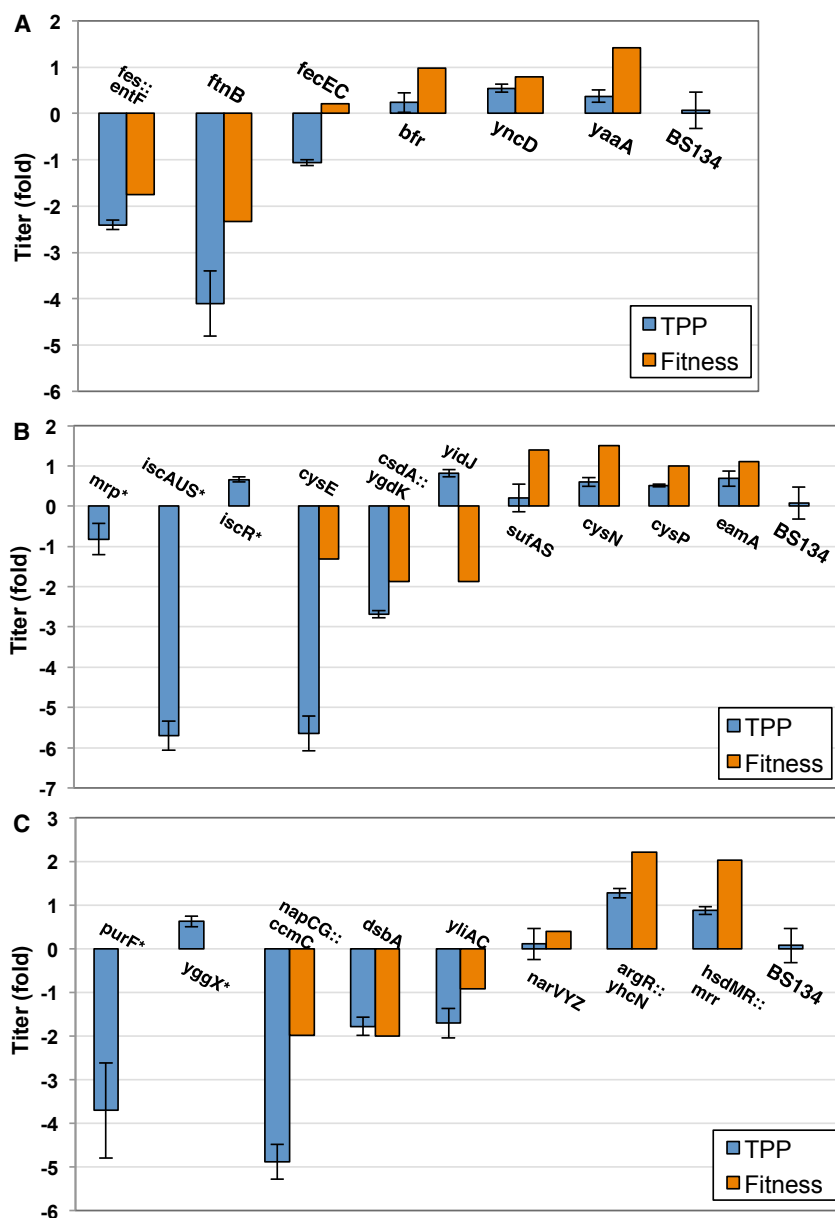
of the OxyR-SoxR and, possibly, Fur regulons, which could be responsible for the distinct effects on iron uptake and sequestration (Figures 3Bi and 3Biv) due to negative feedback (Seo et al., 2014).

### Population Dynamics Effectively Identify Metabolic Targets

We set out to examine whether population-based screening could be used to reliably identify metabolic targets by coupling genetic modification to metabolism. We measured the extracellular titers of TPP produced de novo from the plasmid pTHid in 24 genomic deletions of *E. coli* MG1655 BS134 strain selected from the characterized Fe-S biogenesis network (Figure 3B). For comparison with the newly identified genetic targets, we selected the following five loci previously shown to affect thiamine biosynthesis: *mrp* (Boyd et al., 2008), *purF* (Frodyman et al., 2000), *iscAU* (Agar et al., 2000), *iscR* (Giel et al., 2013), and *yggX* (Grainick and Downs, 2003). We expected to see reductions in biosynthetic output for  $\Delta mrp$ ,  $\Delta purF$ , and  $\Delta iscAUS$  and an increase for  $\Delta iscR$ , a regulator that inhibits transcription from the *iscAUS* operon (Giel et al., 2013). We created 14 single-gene knockouts and ten multi-gene locus lesions (Table S3). For the latter, we expected to observe an additive effect of the genes on TPP output, because their products did not appear to be part of enzyme cascades (Table S4).

We observed a strong correlation between the shift in fitness of a gene knockout in the multiplexed competitive growth assays and the extracellular TPP titer of the isolated gene knockout (correlation: 0.89, excluding the five test loci). Iron supply and availability strongly contributed to TPP output. Disruption of iron storage in ferritin and mobilization from enterobactin completely disrupted the TPP titer ( $\Delta ftnB$  and  $\Delta fes$ -*entF*; Figure 4A). In contrast, knockouts of two genes induced by Fur-OxyR regulons—*bfr* and *yaaA*—whose mutants showed fitness improvements in competitive assays (Figure 3Biv), yielded higher extracellular TPP titers ( $\Delta bfr$ :  $+0.3 \pm 0.2$ ,  $\Delta yaaA$ :  $+0.4 \pm 0.1$ , fold difference from mean titer; see also Figure S5), compared to the average titer across all knockouts and wild-type.

We then investigated pathways involved in S-incorporation into Fe-S in *E. coli*, including free cytosolic cysteine. TPP output decreased, as expected from previous reports (Agar et al., 2000; Boyd et al., 2008; Giel et al., 2013), in the two test loci  $\Delta mrp$  and  $\Delta iscAUS$  (fold difference:  $-0.8 \pm 0.04$  and undetected, respectively). In contrast, knockout of the repressor *iscR* resulted in a higher TPP titer ( $+0.7 \pm 0.06$ ; Figure 4B). We found that disrupting cysteine formation and the mobilization of its sulfur by the cellular sulfur transfer system CSD significantly decreased the TPP titer compared to the mean knockout effect ( $\Delta cysE$ : undetected;  $\Delta csdA$ -*ygdK*,  $-2.7 \pm 0.09$ ), which is in agreement with the reduced fitness of mutants in its components (Figure 3Bii). We previously found that mutants in the *E. coli* SUF sulfur transfer system and in various proteins involved in sulfate assimilation and export exhibited improved growth during TPP selection (Figure 3Bii). The TPP output of single-locus knockouts agreed with these observations: the average extracellular TPP titers were higher in these knockouts



**Figure 4. TPP Quantification in a Set of Genetic Targets**

(A–C) Extracellular titers for individual locus deletions compared to the average titers across all strains (fold difference; for absolute values, see Figure S5). Strains are grouped in three different sub-networks involved in Fe-S biogenesis: iron supply (A), sulfur supply (B), and oxidative regulation (C). Asterisk indicates test KO; fitness bar of multi-gene loci shows average. Error bars indicate SD; n = 3–5. See also Figure S5 and Table S4.

significantly reduced TPP titers:  $\Delta dsbA$ , which is involved in disulfide bond formation (Depuydt et al., 2009) ( $-1.8 \pm 0.2$ ),  $\Delta yliAC$  ( $-1.7 \pm 0.3$ ), and  $\Delta napCG$  (undetected). Mutants in these loci showed reduced fitness during TPP production, whereas those in the SoxR-OxyR regulons had higher fitness. These observations suggest a role for the SoxR-OxyR regulon in diminishing thiamine biosynthesis. Indeed, knockouts of two downstream genes activated by OxyR—the peroxide-response genes *yhcN* (Lee et al., 2010) and *yggX* (Gralnick and Downs, 2003)—had higher extracellular TPP titers (fold difference:  $\Delta yggX$ ,  $+0.6 \pm 0.1$ ;  $\Delta argR\text{-}yhcN$ ,  $+1.3 \pm 0.1$ ) (Figure 4C), in agreement with the shifts in fitness of their pooled mutants (Figure 3Biii; Table S3).

### Tuning of Central Fe-S Control

We examined the role of specific regulators of the complex *E. coli* SoxR-OxyR-Fur regulons during TPP production. We quantified the extracellular TPP titer obtained with  $\Delta soxR$ ,  $\Delta oxyR$ , and  $\Delta fur$  single-gene knockouts introduced in *E. coli* strain TOP10, which lacks DNA restriction systems, and the production strain BS134. We found that, compared to the background strains, the deletion of SoxR did not lead to

increased TPP titers, but the deletion of both OxyR and Fur dramatically increased pathway output in both TOP10 and BS134 (Figures 5A and 5B, respectively). This result pinpointed the OxyR-Fur regulon, which directs iron homeostasis (Seo et al., 2014), as the central cellular response to TPP production.

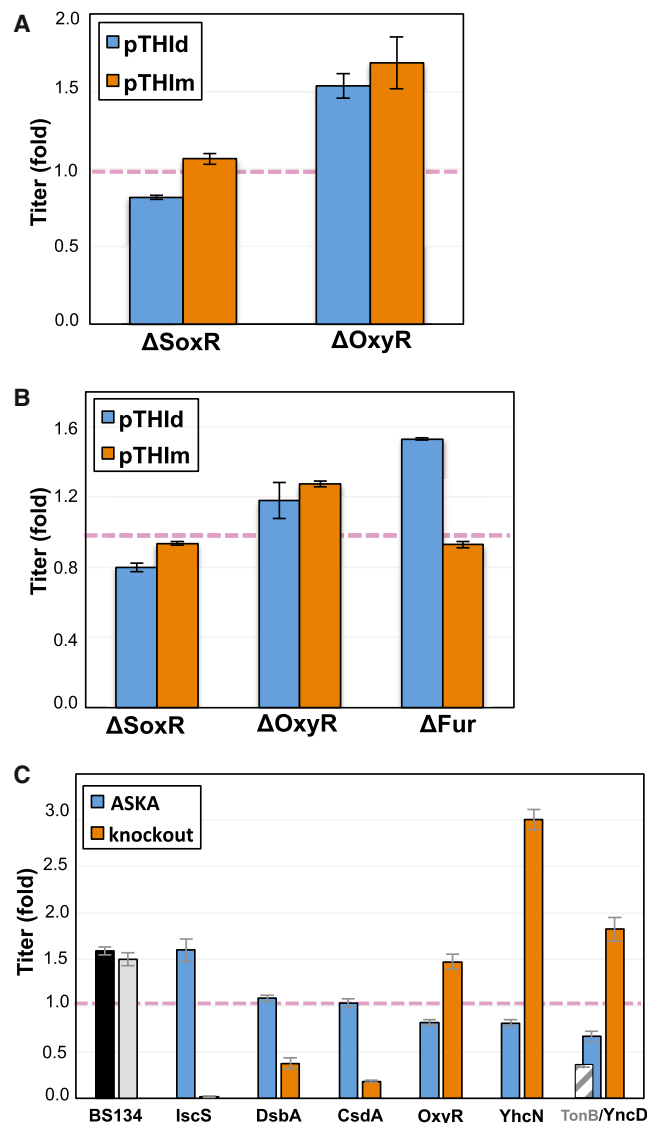
In order to determine whether this response and Fe-S cluster formation could be tuned via the identified genetic targets, we overexpressed the following genes as GFP fusions (Kitagawa et al., 2005): *iscS*, *dsbA*, *csdA*, *oxyR*, *yhcN*, *yncD*, and *tonB*, in the same strain BS134 used for knockouts. Compared to knockouts, TPP titers resulting from gene overexpression showed an opposite shift. In particular, the titer increased during overexpression of *iscS* and *dsbA*, whereas it decreased during overexpression of *oxyR*, *yhcN*, and *yncD* (Figure 5C).

compared to other knockouts and wild-type (fold difference:  $\Delta sufAS$ ,  $+0.4 \pm 0.2$ ;  $\Delta cysN$ ,  $+0.6 \pm 0.1$ ;  $\Delta cysP$ ,  $+0.5 \pm 0.04$ ;  $\Delta eamA$ ,  $+0.7 \pm 0.2$ ) (Figure 4B).

The remaining tested loci encoded proteins involved in redox control of a number of cellular functions, particularly iron homeostasis, which is tightly regulated by the SoxR-OxyR-Fur regulons (Jang and Imlay, 2010; Seo et al., 2014). The OxyR regulon is activated by an environment that favors disulfide bonds, in which glutathione plays a key role (Georgiou, 2002), and by nitric oxide via S-nitrosylation (Seth et al., 2012). Deletion of the *narVYZ* locus, for which a mild fitness effect was calculated ( $\Delta F$ ,  $+0.4$ ), showed TPP titers close to the knockout average ( $\Delta narVYZ$ ,  $0.10 \pm 0.3$ ) (Figure 4C). In contrast, we found that knockouts of the three predicted activators of the OxyR regulon exhibited

increased TPP titers, but the deletion of both OxyR and Fur dramatically increased pathway output in both TOP10 and BS134 (Figures 5A and 5B, respectively). This result pinpointed the OxyR-Fur regulon, which directs iron homeostasis (Seo et al., 2014), as the central cellular response to TPP production.





**Figure 5. Reverse-Engineering OxyR and Fe-S Regulation**

Extracellular titers obtained by engineering central Fe-S cluster formation and the cellular stress response.

(A and B) TPP titer measured in knockouts of central oxidative stress response in *E. coli* TOP10 (A) and BS134 (B) strains bearing pTHld and pTHlm production plasmids (blue and orange, respectively; pink dashed line indicates background strain;  $n = 6-8$ ).

(C) Extracellular TPP titer quantified in *E. coli* BS134 overexpressing several identified genetic targets in Fe-S cluster metabolism and oxidative response (blue bars) compared to their deletion (knockouts, orange bars). Pink dashed line indicates average production;  $n = 6-8$ .

Error bars indicate SD.

## DISCUSSION

The study shows that, by coupling the growth of genome-wide bar-coded Tn mutants to a biosynthetic product, it is possible to identify product-specific metabolic and regulatory bottlenecks. We show that disruptions in the assimilation of certain

forms of iron and sulfur led unexpectedly to improved fitness upon selection for TPP production (Figures 3Bi and 3Bii), a result that was supported by measuring the extracellular TPP titer from individual gene knockouts (Figure 4). Further, overexpression of *tonB*, involved in the uptake of iron-siderophore chelates, led to a significant decrease in TPP titer (Figure 5C). We included in our validation genes that are poorly characterized or of only predicted function, which constitute the so-called Y genes within the *E. coli* genome, and that have strong fitness phenotypes with regard to TPP production. Two genes in particular have predicted function in siderophore uptake (*yncD*, a TonB-dependent receptor) and aryl-sulfate assimilation (*yidJ*, a predicted sulfatase). Both *yncD* and *yidJ* knockouts exhibited TPP titers 20%–25% higher than that of wild-type cells (Figure 4), further confirming that certain assimilatory activities may play a role in inhibiting TPP biosynthesis, possibly through the regulator Fur.

Applied to TPP biosynthesis, genetic-metabolic coupling uncovered that the regulation of Fe-S homeostasis mediated by OxyR-Fur regulons diminishes biosynthetic output by depleting or sequestering cellular Fe-S clusters. In fact, Fe-S clusters are critical for the function of TPP pathway enzymes ThiC and ThiH. The effects of changing global regulators are notoriously idiosyncratic. Nevertheless, TPP-coupled fitness data of mutants in the regulons could be reproduced in different *E. coli* strains and production plasmids (Figures 5A and 5B). Although mutants were not screened for fitness against wild-type cells at the population level, nine of ten selected high-fitness mutants, when introduced in wild-type, exhibited an extracellular TPP titer 0.2- to 2-fold higher (with the exception of *narVYZ*), while, excluding *yidJ*, all other low-fitness mutants exhibited significantly lower titers (Figure 4). We also demonstrate that population-wide mutant fitness data could serve as a basis for tuning a heterologous pathway through overexpression of genetic targets from multi-copy plasmids (Figure 5C).

The characterization of how important parts of the *E. coli* stress response and nutrient assimilation affect a complex, heterologous pathway and the identification of poorly characterized genes—which, today, still represent ~30% of the *E. coli* genome, contributing to this response—demonstrate the power of genetic-metabolic coupling for the identification of target genes for the metabolic engineering of a specific biosynthetic product.

## EXPERIMENTAL PROCEDURES

### Strains and Plasmids

Plasmid DNA, PCR products, and gel extractions were prepared with appropriate kits (QIAGEN). Synthetic oligonucleotides were purchased from Integrated DNA Technologies (IDT). In-frame GFP fusions of selected genes for expression from multi-copy plasmids were obtained from the ASKA library (Kitagawa et al., 2005). Single-locus deletions were constructed by using lambda red recombineering. The thiamine selection system (pTPP\_Bios) was described in a previous study from our laboratory (Genee et al., 2016). The TPP biosynthetic pathway was assembled from native *E. coli* MG1655 genes via PCR amplification. A detailed description of the molecular tools is provided in the Supplemental Information.

### Analytical Quantification of TPP

Extracellular TPP concentration was measured using a modified thiochrome high-performance liquid chromatography (HPLC) assay procedure. Briefly,

fresh colonies carrying either pTHld or pTHlm were inoculated into 1 mL MOPS-rich media lacking thiamine and containing 30  $\mu$ g/mL chloramphenicol and 50  $\mu$ g/mL spectinomycin. Cultivations in deep-well plates were brought twice to saturation in a 48-hr period (Supplemental Information).

### Construction of Tn Mutants and TnSeq and BarSeq protocols

For construction of tagged Tn mutants of *E. coli* MG1655 and bioinformatic analysis of NGS data, we relied on reported methodology and tools (Wetmore et al., 2015). BarSeq reads are converted into a table reporting the number of times each bar code is observed using a custom Perl script (MultiCodes.pl). Given a table of bar codes, in which they map in the genome and their counts in each sample, strain and gene fitness values are estimated with a custom R script (FEBA.R) (Supplemental Information). The bioinformatic tools can be accessed at <https://bitbucket.org/berkeleylab/feba>.

### Calculation of Gene Fitness

Gene fitness is the weighted average of Tn mutant strain fitness within the gene; that is, strains with more reads have less noisy fitness estimates and are weighted more highly. In more detail, we first selected a subset of strains and genes that have adequate coverage in the time-zero samples (at least two reads per strain and 20 reads per gene are considered adequate). Only strains that lie within the central 10% to 90% of a gene coding region are considered. Then, for each sample:

$$\text{Strain fitness} = \text{Log}_2 \left( n_{\text{After}} + \sqrt{\psi} \right) - \text{Log}_2 \left( n_0 + \sqrt{\frac{1}{\psi}} \right)$$

where  $\psi$  is a “pseudocount” to control for very noisy estimates when read counts are very low. (Further details can be found in the Supplemental Information and in Wetmore et al., 2015.)

### SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, five figures, and four tables and can be found with this article online at <http://dx.doi.org/10.1016/j.celrep.2017.07.015>.

### AUTHOR CONTRIBUTIONS

Conceptualization, S.C. and M.O.A.S.; Methodology, S.C. and M.O.A.S.; Investigation, S.C. and F.G.T.; Writing – Original Draft, S.C.; Writing – Review & Editing, S.C. and M.O.A.S.; Funding Acquisition, M.O.A.S.

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